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Match Selection and Refinement for Highly Accurate Two-View Structure from Motion

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Abstract. We present an approach to enhance the accuracy of structure from motion (SfM) in the two-view case. We first answer the question: “fewer data with higher accuracy, or more data with less accuracy?” For this, we establish a relation between SfM errors and a function of the number of matches and their epipolar errors. Using an accuracy estimator of individual matches, we then propose a method to select a subset of matches that has a good quality vs. quantity compromise. We also propose a variant of least squares matching to refine match locations based on a focused grid and a multi-scale exploration. Experiments show that both selection and refinement contribute independently to a better accuracy. Their combination reduces errors by a factor of 1.1 to 2.0 for rotation, and 1.6 to 3.8 for translation.

1 Introduction

3D reconstructions from pictures are increasingly being used to model real scenes or objects. For some applications such as video games or virtual film sets, capturing the general shape and appearance is enough. The reconstruction method does not have to be particularly accurate. However, in industrial settings, where 3D models are used for measurement, accuracy is crucial. Moreover, even for less demanding tasks, accurate reconstruction reduces the quantity of required images, thus reducing the costs and increasing the applicability. Better estimates also lessen the impact of outliers.

In this paper, we propose a method to greatly enhance the accuracy of two-view structure from motion (SfM), i.e., the estimation of the camera poses (positions and orientations) and of the basic structure of the scene (3D point cloud). As 3D reconstruction methods strongly rely on the quality of the estimated calibration, this is a crucial initial step. In most cases, being wrong at calibration time cannot be recovered later.

Match selection. Most SfM approaches are based on the detection and matching of interest points (features) in image pairs [11]. Given point matches between two images, we can estimate a fundamental matrix F relating them. If internal calibration parameters are known (calibration matrix K), this also provides an estimate of the camera motion (rotation R , translation t) and 3D position of matched points. As feature detection and matching is not perfect, two main things can go wrong in the SfM process: the matches can be either incorrect

or inaccurate. There is actually a grey area between these notions: incorrect matches reduce SfM accuracy, sometimes to the point of making it fail, while inaccurate matches are considered as good enough for calibrating, even though they also degrade SfM accuracy. Incorrect matches are generally dealt with using RANSAC [9] or one of its numerous variants [7, 8, 19, 26]. It separates “good matches” (inliers) from “wrong matches” (outliers), trying to find the largest consensus on an estimated fundamental matrix, using a threshold (fixed or adaptive) to assess consistency. While this robust selection method can eliminate many outliers, a number of false positives can remain among the selected inliers because the rejection criterion is mostly based on the distance to epipolar lines, which provides a necessary but not sufficient condition (ambiguity along lines).

Compromises at two different levels impact SfM accuracy. First of all, statistically, the more matches to calculate the fundamental matrix, the more accurate the estimation. A first compromise thus concerns the RANSAC selection criterion: if it is too permissive, matches considered as inliers are more numerous but are also more likely to be contaminated by wrong matches, and accuracy drops; if the criterion is too strict, there are too few inliers to get a good accuracy. The second compromise concerns the accuracy heterogeneity of individual inliers: keeping only the most accurate inliers can naturally improve SfM accuracy; but it can also degrade it as the estimation is based on fewer points. The first compromise has indirectly been widely studied: people try to select as many good matches as possible, while excluding as many wrong matches as possible. But the second compromise, quality vs. quantity, has been poorly addressed. This paper presents an original method to find a good balance between the number of inliers to consider for SfM estimation and their expected accuracy.

Match refinement. Another way to obtain a better SfM accuracy is to improve the accuracy of feature detection and matching.

Due to differences in imaging conditions, in particular changes of viewpoint or illumination, a salient point or region detected in one image is not detected in the other image at the exact same location. The most popular features are by design only invariant (at most) to affine transformation (e.g., Harris-affine [18], MSER [17], ASIFT [20]), or to small affine transformation (e.g., SIFT [16]). But they are not invariant under perspective transformation, which is enough to offset most detections. Methods to add some perspective invariance to existing feature detectors have been proposed, but they require full 3D information (depth map or mesh) [13, 28], which is computationally expensive or requires more than just image data; others are suited to specific classes of scenes only, mostly urban environments, as they strongly rely on the presence of vanishing points and large planar surfaces [2, 5]. Besides, they have been designed to improve the repeatability of feature detection and matching, which is generally measured using a threshold on the relative overlap of corresponding regions [18], not in terms of the closest distance between feature centers.

Traditionally, two detected feature points can nonetheless be considered as matching although their position in the images does not correspond exactly to the same 3D point in the observed scene. For a number of tasks, being close is

enough. But for highly accurate calibration, it is not satisfactory. In fact, we do not care whether a specific 3D point is accurately identified in both images, such as the very tip of a corner. What we need is possibly arbitrary pairs of points in the images as long as they correspond to extremely close 3D points. In this sense, feature detection and matching is just a way for us to identify corresponding regions rather than corresponding points: their center generally corresponds only to close but different 3D points. Our match refinement only uses them as initial estimates to find a neighboring pair of points that is likely to correspond to closer 3D points because they have a better photometric consistency (assuming an unknown affine transformation).

Finely relating image regions can be addressed with optical flow methods [3]. But they are not well adapted here because they suppose small variations, both in viewpoint (very small baseline, quasi-affine transformation) and in illumination (controlled light scenes). Refining the position of image regions to overlap them better has been studied in the photogrammetry community. One of the most popular methods is adaptive least squares matching (LSM), that tries simultaneously to find radiometric and geometric corrections to best fit two images patches [10]. The most complex geometric correction generally considered in this framework is affine transformation, because projective transformations are assumed to be sufficiently approximated by an affinity. We present here an improvement of LSM based on a focused irregular grid and made robust with coarse-to-fine exploration. We show that it outperforms affine correction.

Our contributions follow the structure of the paper: We establish an empirical statistical relationship between the inaccuracy of matches, their number, and various indicators of SfM inaccuracy (Section 2). We describe an original method that exploits this relationship to select matches that are likely to improve SfM accuracy (Section 3). We present a novel method to locally refine the position of matches to improve their accuracy (Section 4). We show on extensive experiments that both methods improve substantially the accuracy of structure from motion, and even more when combined (Section 5). Section 6 concludes.

2 Statistical Behavior of SfM Errors

2.1 Theoretical Results

We consider a pair of images I, I' , obtained by cameras C, C' with 3×4 projection matrices P, P' and 3×3 calibration matrices K, K' . We also consider a set of matches M between I and I' , i.e., pairs of points $m = (\mathbf{x}, \mathbf{x}')$ where \mathbf{x} is the projection of a 3D point X on I , i.e., $\mathbf{x} = PX$ in homogeneous coordinates, and where \mathbf{x}' is a point in I' considered as matching with \mathbf{x} , possibly with some inaccuracy. In the general case, a fundamental matrix F_M between I and I' can be estimated from matches M , and F_M may in turn be used with K, K' to estimate projection matrices P_M, P'_M on I, I' . The resulting reprojection error of \mathbf{x} in I' , i.e., the discrepancy in I' between the exact reprojection of X by P' and the estimated reprojection of X by P'_M is the distance $e_{2D}(M, m) = d(P'X, P'_MX)$.

In case images I and I' are related by homography H , and considering matching points \mathbf{x}' as possibly inaccurate measurements of reprojected points $P'X = H\mathbf{x}$ in I' , Hartley and Zisserman [11, §5.1.3, Eq.(5.5)] show that, if these measurements are subject to independent Gaussian noise with standard deviation $\sigma_{2D}(M)$, then the estimation error $e_{2D}(M)$ of reprojected points in I' by the estimated homography H_M , or equivalently via P'_M , is:

$$e_{2D}(M) = \mathbb{E}_{m \in M} [e_{2D}(M, m)^2 / |M|]^{1/2} = 2\sigma_{2D}(M) / \sqrt{|M|}. \quad (1)$$

Dividing the estimation error by 2 thus requires 4 times as many matches, or matches with location error divided by 2. This bound is optimal (assuming no other errors such as distortion), and achieved for the Maximum Likelihood Estimator (MLE). Finding a similar bound for the fundamental matrix is impractical because it is a non convex problem in very high dimension. We do not try to solve it, but we draw inspiration of the MLE bound in what follows.

Another reading of (1) is that if we can find a subset $M_{\text{sub}} \subset M$ such that matching points \mathbf{x}' in M_{sub} are subject to independent Gaussian noise with standard deviation $\sigma_{2D}(M_{\text{sub}}) < \sigma_{2D}(M)$ compared to their expected location $H\mathbf{x}$, and if $\sigma_{2D}(M_{\text{sub}}) / \sqrt{|M_{\text{sub}}|} < \sigma_{2D}(M) / \sqrt{|M|}$, then $e_{M_{\text{sub}}} < e_M$, and $H_{M_{\text{sub}}}$ is thus a better estimate of H than H_M . Now if we have a way to evaluate $\sigma_{2D}(M_{\text{sub}})$ for any M_{sub} , the optimal subset M_{sub}^* of matches for estimating H is:

$$M_{\text{sub}}^* = \arg \min_{M_{\text{sub}} \subset M} \sigma_{2D}(M_{\text{sub}}) / \sqrt{|M_{\text{sub}}|}. \quad (2)$$

$H_{M_{\text{sub}}^*}$ minimizes reprojection errors w.r.t. ground truth H (not w.r.t. $H_{M_{\text{sub}}^*}$ that has a trivial solution with any 4 points).

To our knowledge, a similar result is not known for the fundamental matrix. The situation is more complex in this case as estimating F , with 7- or 8-point methods, relies on SVD and/or requires solving complex polynomial systems.

2.2 Empirical Results

As a theoretical result is difficult to obtain, we study empirically the influence of $|M|$ and $\sigma_{2D}(M)$ on SfM accuracy. Using a collection of images with accurate ground-truth calibration, presenting various feature distributions, we measure:

- F_M is the fundamental matrix estimated from M using ORSA (a RANSAC variant) and iterative re-weighted least squares (IRLS) [19].
- $e_F(M)$ is the root mean square error (RMSE) of the distance $e_F(M, m)$ of \mathbf{x}' to the F_M -epipolar line of \mathbf{x} in I' , for $m=(\mathbf{x}, \mathbf{x}') \in M$.
- $e_R(M) = \angle R_{\text{gt}} R_M^{-1}$ is the angle between the ground-truth rotation R_{gt} and its estimate R_M based on M .
- $e_t(M) = \angle(t_{\text{gt}}, t_M)$ is the angle between the ground-truth translation direction t_{gt} and its estimate t_M .
- $e_{3D}(M, R, t)$ is the RMSE of the distance of the 3D point \hat{X} triangulated from \mathbf{x}, \mathbf{x}' using a given rotation and translation R, t , to the ground-truth 3D point X , for $m=(\mathbf{x}, \mathbf{x}') \in M$. We also define $e_{3D}(M) = e_{3D}(M, R_M, t_M)$.

Realistic, semi-synthetic dataset. Estimating SfM errors requires a ground truth for both calibration and matched points. While accurate camera calibrations can be determined using LiDAR data [23], it is difficult to construct a significant number of accurate ground-truth point matches. For this, we resort to semi-synthetic ground-truth datasets: the images, the camera poses and the distribution of matching points are real, but the actual point locations are adjusted to make sure they are error-free.

Concretely, given a pair of images I, I' with known calibration $P_{\text{gt}}, P'_{\text{gt}}$, we detect and match SIFT feature points in each image. We use a descriptor distance ratio to next best match at most 0.8, which is the standard setting [16]. As these matches may still contain outliers, we first clean them using the K-VLD method of Liu and Marlet [15], that eliminates many false matches, including near the epipolar lines, and then using ORSA, an adaptive state-of-the-art variant of RANSAC by Moisan and Stival [19], known for its robustness in practical SfM systems [21]. It results in an almost outlier-free set of matches \tilde{M} . Treating them as inliers, for each match $\tilde{m} = (\tilde{\mathbf{x}}, \tilde{\mathbf{x}}') \in \tilde{M}$, we construct a 3D point X by triangulation using ground truth calibration $P_{\text{gt}}, P'_{\text{gt}}$, and reproject it onto images I, I' as new 2D points $(\mathbf{x}, \mathbf{x}') = m$. The resulting set of matches M_{gt} yields a perfect ground truth that is realistic in terms of feature distribution in images (location and number) and in space.

We then add noise by randomly moving in image I' the matched points \mathbf{x}' , using an isotropic Gaussian distribution with given standard deviation σ_{2D} . This asymmetric setting reproduces the theoretical hypothesis mentioned in Section 2.1. (Adding noise to points in both images experimentally leads to almost identical results, scaled by a constant.) To also conform to this hypothesis, the noise is independent of the characteristics of the features that originated the synthetic points, such as scale. Moreover, we add variation to the number of matches by randomly selecting only a given ratio r . This defines new sets $M = M(\sigma_{2D}, r)$.

In our experiments, we use Strecha et al.'s dataset [23]. It consists of 6 groups of 8 to 30 images with both internal and external accurate ground-truth calibration. We consider all pairs of consecutive images in all image groups, in which we detect and match SIFT features. The number of matches typically varies between 300 and 6000. For each image pair, we consider discrete ratios of matches $r = k^2/100$ with $k = 4, \dots, 10$ (thus different point configurations), and discrete deviation $\sigma_{2D} = 0.2 + 0.3k$ with $k = 0, \dots, 6$ (in pixels). For each combination of r and σ_{2D} , we sample 50 noisy variants of the data, estimate their SfM accuracy, and average the corresponding error measures by quadratic mean (RMS).

Analysis. Adding noise σ_{2D} and ratio r as $M = M(\sigma_{2D}, r)$ has a direct impact on epipolar error $e_F(M)$ and on rotation and translation errors e_R, e_t . It has an impact in turn on the error e_{3D} of estimated 3D points, which depends both on noisy matches M and estimated calibration R_M, t_M .

We observe that e_{3D} , e_R and e_t all are highly correlated to $N = |M|$ and σ_{2D} : although there are some variations, we notice experimentally that $\log e_{3D}$, $\log e_R$ and $\log e_t$ are more or less linear with respect to $\log N$ when σ_{2D} is fixed, with some slope α depending on the image pair, and more or less linear with $\log \sigma_{2D}$.

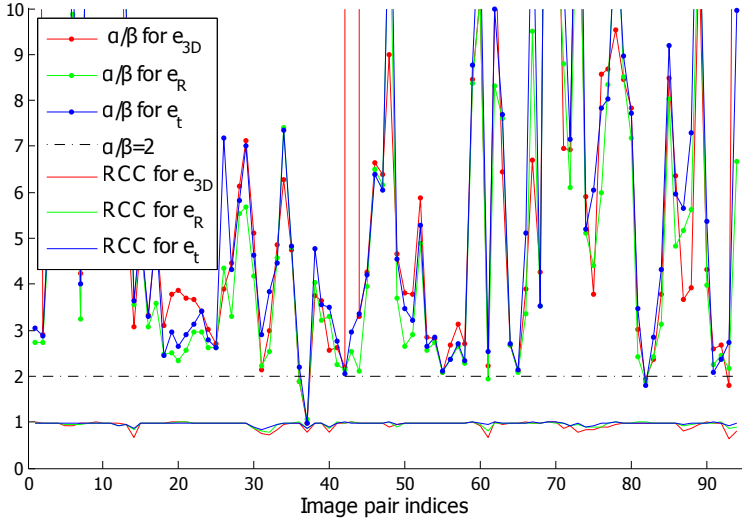


Fig. 1. Dotted curves: estimated α/β for different image pairs (the order is irrelevant). Plain curves: regression correlation coefficient between e_R , e_t , or e_{3D} , and $\sigma_{2D}^\alpha/N^\beta$.

when N is fixed (but not the configuration), with some slope $-\beta$ also depending on the image pair. It is confirmed by computing the regression correlation coefficient (RCC) of e_{3D} , e_R , e_t with $\sigma_{2D}^\alpha/N^\beta$, which is in general very close to 1, as can be seen in Figure 1 (bottom 3 curves, plotted on the same diagram).

Besides, we also found empirically that σ_{2D} is more or less proportional to e_F , not only to the exact epipolar error. We thus hypothesize the relation:

$$e_R, e_t, e_{3D} \propto \frac{\sigma_{2D}^\alpha}{N^\beta} \propto \frac{e_F^\alpha}{N^\beta} \quad (3)$$

With a fixed distribution of points, we should have $\alpha = 1$ for small errors. Still with a fixed configuration, but duplicating all matches, the covariance matrix of estimated parameters is halved; we should thus have $\beta = 0.5$. This is consistent with equation (1). However, it does not hold when point configuration varies. Experimentally, α and β can vary significantly depending on images pairs and match sampling. In our semi-synthetic dataset, β varies between 0.2 and 1.5. Yet, assuming relation (3), knowing α/β is sufficient to compare errors for a given image pair:

$$\frac{e_F^\alpha}{N^\beta} < \frac{e_F'^\alpha}{N'^\beta} \Leftrightarrow \frac{e_F^{\alpha/\beta}}{N} < \frac{e_F'^{\alpha/\beta}}{N'} \quad (4)$$

The situation where all matched points are treated as inliners and contribute to estimating F amounts to preferring the largest N (smallest $1/N$) independently of e_F , i.e., to $\alpha/\beta = 0$. On the contrary, the larger α/β , the more aggressively low-accuracy features should be discarded. As can be seen in Figure 1,

$$\alpha/\beta \geq 2 \text{ almost consistently.} \quad (5)$$

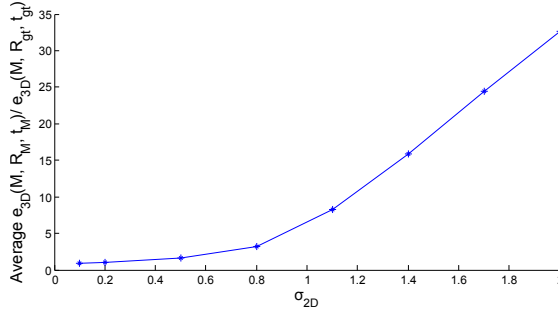


Fig. 2. Amplification of 3D reconstruction error when σ_{2D} grows

Figure 2 compares the accuracy of reconstructing 3D points from noisy image points with ground-truth R_{gt}, t_{gt} vs estimated R_M, t_M (average on image pairs). The bigger σ_{2D} , the more calibration errors amplify reconstruction errors.

3 Match Selection to Improve Accuracy

To improve accuracy, we estimate SfM using a selected subset of good matches.

Cleaning up input matches. Although we use IRLS for estimating F , the level of accuracy we target may be sensitive to outliers remaining after RANSAC. We thus try to eliminate outliers from the set of input matches. We have to do it without introducing the bias of an early approximate calibration estimation, which would be the case if we were to first filter the matches using RANSAC. For this reason, we first clean up the matches using the K-VLD method [15]. Based on semi-local geometric and photometric consistency, it eliminates many outliers without any calibration assumption. Running ORSA afterwards on the resulting set of matches M typically only removes on the order of 10% of matches with a found threshold of less than 2-pixel error for estimating F .

Comparing subsets of matches. The SfM errors we want to reduce are $e_R(M), e_t(M), e_{3D}(M)$. But what can be easily measured given a pair of images and a set of matches M is just $e_F(M)$. However, as indicated by Eq. (3) and (4), $e_R(M), e_t(M), e_{3D}(M)$ vary monotonically with $e_F(M)^{\alpha/\beta}/|M|$. The basic idea of match selection is to use only a subset of matches $M_{sub} \subset M$ as soon as:

$$\frac{e_F(M_{sub})^{\alpha/\beta}}{|M_{sub}|} < \frac{e_F(M)^{\alpha/\beta}}{|M|} \quad (6)$$

However, α/β is a priori unknown for an arbitrary image pair. Moreover, we want to improve SfM without taking the risk to degrade it. What we need is a sufficient condition that reducing the number of matches will probably improve accuracy but most certainly will not reduce it. For this, we look for a possible value $\gamma \geq 0$ such that, for any image pair, any set of matches M with corresponding α, β parameters, and any subset of matches $M_{sub} \subset M$,

$$\frac{e_F(M_{sub})^\gamma}{|M_{sub}|} < \frac{e_F(M)^\gamma}{|M|} \Rightarrow \frac{e_F(M_{sub})^{\alpha/\beta}}{|M_{sub}|} < \frac{e_F(M)^{\alpha/\beta}}{|M|} \quad (7)$$

We can then choose this optimal subset of matches M_{sub}^* for estimating F :

$$M_{\text{sub}}^* = \arg \min_{M_{\text{sub}} \subset M} \frac{e_F(M_{\text{sub}})^\gamma}{|M_{\text{sub}}|} \quad (8)$$

The fundamental $F_{M_{\text{sub}}^*}$ minimizes reprojection errors w.r.t. ground truth F_{gt} .

Noting that $(e_F(M_{\text{sub}})/e_F(M))^\gamma < |M_{\text{sub}}|/|M| < 1$ and hypothesizing (5), we can choose $\gamma = 2$ because then $(e_F(M_{\text{sub}})/e_F(M))^{\alpha/\beta} < (e_F(M_{\text{sub}})/e_F(M))^\gamma$, ensuring condition (7). Parameter γ is chosen as a safe empirical lower bound, not an average value, which is more robust. Still, a general method to treat a specific class of images would be to run experiments as in Section 2.2 and pick a value $\gamma \leq \alpha/\beta$. Without loss of generality, we assume $\gamma = 2$ in the following.

Exploring subsets of matches. The difficulty to find M_{sub}^* is to explore $M_{\text{sub}} \subset M$, as there are too many such subsets ($2^{|M|}$). We propose to evaluate just a fraction of them, that has the most chances to lead to smaller ratios $e_F(M_{\text{sub}})^2/|M_{\text{sub}}|$. For this, we rank the matches in M and use this ordering to explore only subsets of top-rank matches. More precisely, we look for a ranking function $\phi: M \rightarrow \mathbb{R}$ to order the matches into a sequence $(m_i)_{1 \leq i \leq |M|}$ such that $i < j \Rightarrow \phi(m_i) < \phi(m_j)$, and consider $M_{\text{sub}}(N) = \{m_i \mid 1 \leq i \leq N\}$. If the ranking function ϕ is highly correlated to the reprojection errors $e_{2D}(M, m)$, and hence to the epipolar errors $e_F(M, m)$, then

$$\begin{aligned} \min_{M_{\text{sub}} \subset M} \frac{e_F(M_{\text{sub}})^2}{|M_{\text{sub}}|} &= \min_{N \leq |M|} \frac{1}{N} \min_{\substack{M_{\text{sub}} \subset M \\ |M_{\text{sub}}|=N}} e_F(M_{\text{sub}})^2 \\ &\approx \min_{N \leq |M|} \frac{1}{N} e_F(M_{\text{sub}}(N))^2 \end{aligned} \quad (9)$$

We may thus resort to:

$$N^* = \arg \min_{N \leq |M|} \frac{e_F(M_{\text{sub}}(N))^2}{N} \quad (10)$$

$$M^* = M_{\text{sub}}(N^*) \quad (11)$$

The number of subsets to explore is then reduced from $2^{|M|}$ to $|M|$, which is still a lot given that M generally contains a few thousands of matches. Note that $e_F(M_{\text{sub}}(N))^2/N$ is not necessarily convex. However, it is in practice “smooth” enough for a reduced exploration of $8 \leq N \leq |M|$ to make sense. In our various experiments, we found it robust and accurate enough to ensure a minimum of 40% of matches in $M_{\text{sub}}(N)$ and to explore fractions of M with a 5% step, i.e., to consider $N = r|M|$ with ratio $r = 0.4 + 0.05k$ and $k = 0, \dots, 12$.

Ranking matches. The choice of a ranking function ϕ varies with the kind of feature. For SIFT, it seems natural to consider the distance between feature descriptors $d(\text{desc}(\mathbf{x}), \text{desc}(\mathbf{x}'))$ as an indicator of feature accuracy. Besides, Tang [25] showed that SIFT subsampling amplifies location error by the feature scale factor $\text{scale}(\mathbf{x})$. This leads us to define the following ranking function:

$$\phi(\mathbf{x}, \mathbf{x}') = \max(\text{scale}(\mathbf{x}), \text{scale}(\mathbf{x}')) d(\text{desc}(\mathbf{x}), \text{desc}(\mathbf{x}')) \quad (12)$$

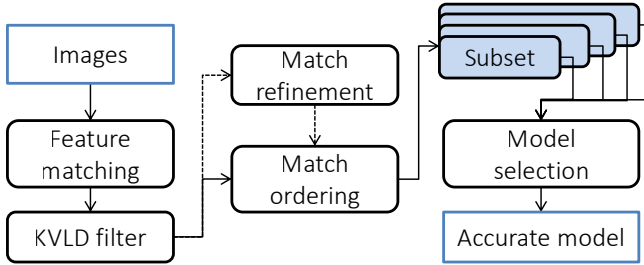


Fig. 3. A global view of our algorithm

Large features thus tend to be ordered last, unless their descriptors match well. Still, although they have a poor accuracy, they are often useful for robustness, which could be a issue if too many of them are discarded. But our use of K-VLD [15] provides enough (if not better) robustness improvement to compensate.

On semi-synthetic data, made with first images from Mikolajczyk et al.’s dataset [18] and after applying an known homography, we found a correlation coefficient of 0.42 between ϕ and σ_{2D} , which proves the relevance of ϕ for ordering M . It outperforms other indicators, such as feature saliency that has a correlation score of 0.02. The Lowe score (ratio of descriptor distance to next best match) has an individual correlation of 0.2, but it does not improve the global correlation when combined with ϕ .

Note that the definition of ϕ relies only on detection scale and on the SIFT *descriptor*, not on the detector. It can thus be used, e.g., for all detectors of Mikolajczyk et al. [18], including SURF, Harris-Affine and MSER. Transposition to other descriptors is direct, but the correlation coefficient should be checked.

Algorithm. Our match selection algorithm is summarized on Figure 3. After feature detection and matching, matches M are cleaned up using K-VLD and ordered using the ranking function ϕ . Subsets M_{sub} of sorted matches are explored to minimize $e_F(M_{\text{sub}})^\gamma / |M_{\text{sub}}|$ and the subset with the lowest value if used to construct the estimated model.

Comparison to related methods. The PROSAC variant of RANSAC also constructs a series of match subsets and iterates first on better ones [7]. However, the target is not accuracy but fast convergence; robustness and precision are similar to RANSAC. Note that our method is not an alternative to RANSAC nor a fundamental matrix estimator, but a complement: a RANSAC variant as well as a fundamental matrix estimator are still needed to compute the calibration and the corresponding epipolar error e_F for the different M_{sub} subsets considered. As a matter of fact, Section 5 shows that our method, combined with different variants of RANSAC, consistently provides much better results.

4 Least Square Focused Matching

We now present an extension of least squares matching (LSM) [10, 22] to better adjust the location of matching features. LSM is based on the hypothesis

that, *locally*, the region around the feature center is mostly planar, so that two matching regions are approximately related by homography, which in turn can be approximated by an affinity if the change of viewpoint is moderate. Besides, image intensity is also considered to possibly vary with an affine transformation.

Finding the affine parameters (both geometric and radiometric) that best map the two regions provides a good estimate of point displacement. This is a non-linear adjustment problem. It can be addressed by an iterative scheme based on a first-order Taylor expansion expressing optical flow constraints: differentiating the affine relation between the two regions, a small change in the affine parameters can be related to a small change in the dissimilarity measure of the regions, to be minimized. Yet, contrary to ordinary optical flow [3], LSM is not restricted to small light changes, small rotation angles and small baselines.

With LSM, we propose two improvements. First, instead of using a regular sampling grid around the features, we use an irregular grid focused on the center of the region to match. Second, we combine it with an image scale traversal to make it more robust to local minima. (We also tried estimating a homography rather than just an affinity, but it did not produce substantial improvements.)

Note that feature detection covariance [4, 12, 24, 29] is irrelevant here. What we do is, given a position p in I for which we know a roughly corresponding position p' in I' , adjust p' so that the regions around p in I and p' in I' correlate better, under some geometric and photometric affinity to estimate. Feature points that match just happen to provide good initial correspondences for the refinement process. This is also widely different from refining the location of features detected as salient [16, 18]. Moreover, refining given matches leads to a better accuracy than refining detections before matching.

Initialization. Given two matching features, we measure the dissimilarity η between the region with the lowest scale s and the region with the highest scale s' after enlarging by a factor of s'/s the sampling grid. We also rotate the grid according to the difference of orientation between the features. This initializes the affinity parameters in the iteration process.

Focused grid. The image dissimilarity measure η in LSM is traditionally based on a regular sampling grid centered on interest points. This assumes a uniform transformation of the whole patches, which is true locally and slowly breaks down when moving away from the center. For this reason, we propose to use a grid focused on the patch center, i.e., denser in the center than in the border, which is additionally weighted by a Gaussian kernel to further concentrate on the center. Concretely, we use a grid whose lines are defined by a geometric progression: its coordinates are $(u, v) = (\text{sign}(i) \frac{\rho^{|i|}-1}{\rho-1}, \text{sign}(j) \frac{\rho^{|j|}-1}{\rho-1})$ for $i, j \in \{-n, \dots, n\}$. A standard Gaussian weight of $\frac{1}{2\pi\sigma^2} \exp(-\frac{u^2+v^2}{2\sigma^2})$ is used on grid nodes (u, v) . Figure 4 illustrates the shape of our grid, with color representing the Gaussian weight. In experiments, we use a spline interpretation of order 5 to get subpixel intensity in a focused grid with $n = 7$, $\sigma = 0.9n$ and $\rho = 1.1$: samples at the grid border are then almost twice as dense (in one direction) as at the center.

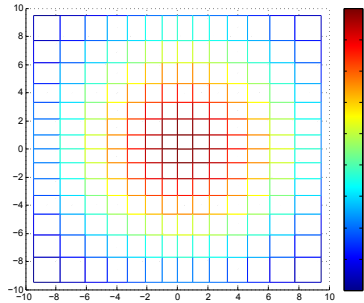


Fig. 4. Focused grid: denser and heavier center (colors represent the Gaussian weight)

Scale exploration. Additionally, rather than directly adjusting the feature positions at their original scale, we perform a coarse-to-fine refinement. We start adjusting point location at a higher scale and progressively refine the location by reducing the scale until we reach the original scale. (An optimal scale is not search as in [14]: the original scale is best for accuracy.) It improves robustness, preventing some refinements to be caught in local minima. For this, we create a pyramid of images similar to the one used in SIFT detection. After convergence of the geometric and photometric parameters at a given scale, we restart with the corresponding location and parameters at the scale below. As a high blur may also cause deviation from the optimal solution, we make sure there is an actual improvement: if the measure of dissimilarity computed with the estimated parameters at the current scale is less than the dissimilarity on the lower scale using just feature scale and orientation, the latter is kept as initial parameters for the refinement at the lower scale. In our experiments, we explore 5 octaves of scale, dividing each octave in 2, i.e., with a geometric progression of ratio $\sqrt{2}$.

Impact of the feature detector. SSD-based refinement is more accurate for regions with high gradients. SIFT does not necessarily detect points in such regions, but its robustness compensates. Besides it tends to find points within objects, where relative intensity is more stable, compared to corners that have strong but less stable gradients because they often correspond to occlusion edges.

Match selection with match refinement. Match selection and match refinement are independent improvements that can be combined, match refinement coming first (see Fig. 3). However, match refinement changes the correlation between the match errors and the indicators of feature accuracy. When combined, the match ranking to create subset candidates (cf. Section 3) has to be changed.

Based on experiments with the same semi-synthetic data as in Section 3, we found that, after match refinement, the dissimilarity measure η using the focused grid has correlates with the actual feature localization error, with score of 0.27. Besides, intuitively, the scaling and shearing of the image, as defined by the affinity estimate A , also has an impact on the quality of matching. Given orthogonal vectors (u, v) in I , we consider the value $\max_{u,v} \frac{|u^T A^T A v|}{|A u| |A v|}$, which is the cosine of the maximum crushing after transformation. It can be shown to be

simply expressed as $\chi = \frac{|\lambda_1 - \lambda_2|}{\lambda_1 + \lambda_2}$, where $\lambda_1, \lambda_2 > 0$ are the eigenvalues of $A^T A$. It has a correlation score of 0.12 with the localization error. By a linear regression over the same semi-synthetic data, we empirically define $\phi(m) = 0.3\eta + 42.6\chi$, which has a correlation of 0.34 with the location error. Note that feature scales no longer correlate with the location error (correlation is only 0.01) and are thus discarded from the ranking function.

5 Experiments

To evaluate our method, we consider some RANSAC variants among those that are considered the most suited for accuracy (as opposed, e.g., to robustness or speed) [6, 27]: RANSAC with iterative re-weighted least squares (IRLS) for final model estimation [27, method S1], RANSAC with M-estimator (MSAC), LO-RANSAC [8], MLESAC [26], and ORSA with IRLS [19]. IRLS tries to minimize the sum of squares of geometric error between points in the right image and the epipolar line of corresponding points in left image. For each of these variants, we compare 4 settings: RANSAC alone, RANSAC preceded by match selection (MS), RANSAC preceded by match refinement (MR) using LSFM, and RANSAC preceded both by match refinement and match selection (MR+MS). A uniform threshold of 3 pixels (distance to epipolar line) is used in the RANSAC variants for outlier rejection, apart from ORSA that chooses the threshold automatically. All the results we provide are averaged over 20 runs.

Only datasets with highly accurate ground-truth calibration can be used for validation. We experimented with the full dataset of Strecha et al. [23], a de facto standard in camera calibration: 6 groups of 8 to 30 images totaling 95 pairs of successive images. For each pair, SIFT feature points are detected and matched with the usual setting [16] (no tweaking as in Sect. 2.2), i.e., a descriptor distance ratio to next best match at most 0.8. We ran the same experiment with the DTU robot dataset [1]. However, as it is huge (about 0.5 To), we only considered 9 of the 60 groups of images, covering various themes (scenes 1, 2, 4, 9, 10, 12, 21, 28, 52), in the reduced format (fewer images, yielding 12 images pairs: 1-12, 12-24, 24-25, 25-26, 26-37, 37-49, 50-57, 57-64, 57-65, 57-94, 64-95, 64-119), with identical illumination condition (number 08 for all tests), but full-size images.

Match selection and refinement. Figure 5 shows the average rotation and translation errors e_R, e_t for each scene of each dataset. Table 1 shows the average results, illustrating both the separate and combined benefits of MS and MR. Gain factors attain 2.0 for rotation and 3.8 for translation. Note that most parameters are learned on other, widely different images [18]; only the lower bound $\gamma = 2$ is defined from feature distribution in [23] and nothing else. Our excellent results on [1, 23] suggests that these parameters make sense for a wide range of images.

Focused matching. We compare our focused matching (LSFM, Section 4) with standard least square matching (LSM). Rather than considering planar scenes and measuring reprojection errors, we directly measure e_R and e_t using the datasets and settings as above. Matches are then refined by the following

Table 1. Average rotation and translation errors: RANSAC alone (raw), + match selection (MS), + match refinement (MR), + both (MR+MS), and gain raw/(MR+MS)

Dataset	Strecha et al. [23]					DTU robot [1]				
e_R (deg $\times 10^{-2}$)	raw	MS	MR	MR+MS	gain	raw	MS	MR	MR+MS	gain
RANSAC	16.4	9.52	10.3	8.87	1.9	26.5	22.3	21.5	21.3	1.2
MSAC	14.1	9.53	8.86	8.43	1.7	21.3	21.7	20.4	20.1	1.1
LO-RANSAC	16.4	9.54	10.3	8.97	1.8	26.8	22.2	21.5	21.3	1.3
MLESAC	15.8	7.81	9.50	7.76	2.0	21.8	22.6	20.8	20.2	1.1
ORSA	12.2	7.24	6.48	6.60	1.9	21.9	21.7	20.8	20.3	1.1
e_t (deg)	raw	MS	MR	MR+MS	gain	raw	MS	MR	MR+MS	gain
RANSAC	1.85	1.09	1.23	1.04	1.8	3.83	2.12	1.81	1.02	3.7
MSAC	1.59	1.08	1.03	0.96	1.6	1.27	1.03	0.93	0.70	1.8
LO-RANSAC	1.83	1.10	1.21	1.05	1.7	3.89	2.14	1.76	1.02	3.8
MLESAC	2.16	0.95	1.09	0.87	2.5	2.02	1.34	1.23	0.77	2.6
ORSA	1.38	0.81	0.68	0.74	1.9	1.22	0.88	0.66	0.66	1.8

Table 2. Match refinement evaluation using LSM, LSM with focused grid, LSM with focused grid and scale exploration (LSFM), gain as improvement of LSFM over LSM

e_R (deg $\times 10^{-2}$)	LSM	LSM + foc. grid	LSFM	gain
Strecha et al. [23]	7.55	6.73	6.48	1.17
DTU robot [1]	20.97	21.22	20.79	1.01
e_t (deg)	LSM	LSM + foc. grid	LSFM	gain
Strecha et al. [23]	0.84	0.72	0.68	1.25
DTU robot [1]	0.77	0.71	0.66	1.16

methods: LSM, LSM with focused grid, and LSM with focused grid and scale exploration (LSFM). We estimate errors after calibrating with ORSA+IRLS, which has the best performance in the above tests (see Table 1). Table 2 shows that, apart from a poor reduction of rotation error in the DTU robot dataset, the LSFM factor gain is 1.16 to 1.25.

6 Conclusion

In this paper we have studied, in the two-view case, the “quality vs. quantity” balance of point matches for structure from motion — a poorly addressed issue in the literature. We have found a correlation between SfM errors and a function of the number of matches and their epipolar errors. Using this relation, we have presented a new method for selecting relevant subsets of points to improve SfM accuracy. We have also proposed an improvement of an existing method to refine match locations. Using extensive experiments involving real data with ground-truth calibration, we have shown that match selection and match refinement independently lead to a major reduction of SfM errors over the best methods

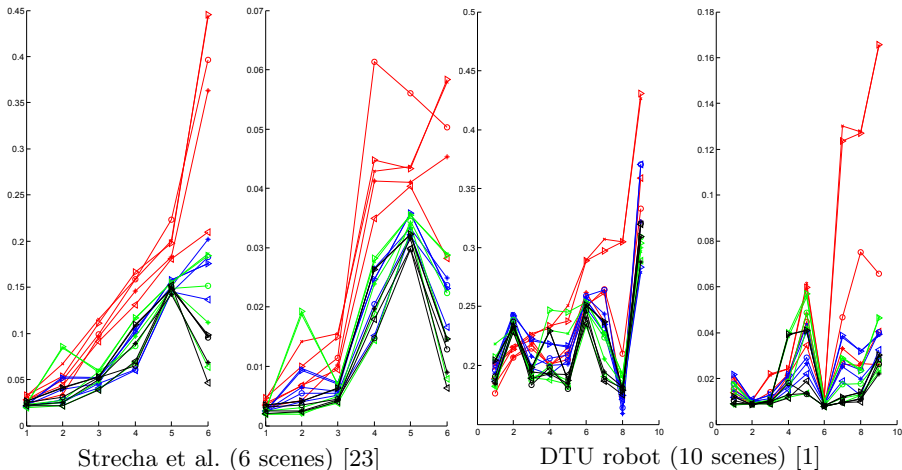


Fig. 5. Average results on the datasets. Left: rotation error e_R . Right: translation error e_t . Color **red**: raw RANSAC; **blue**: with match selection (MS); **green**: with match refinement using LSFM (MR); **black**: with both match selection and match refinement (MR+MS). Line symbol \rightarrow : RANSAC with IRLS; $*$: MSAC; x : LO-RANSAC; o : MLESAC; $<$: ORSA. Scenes are reordered by increasing rotation error of RANSAC

targeted at accuracy. Combining both methods, the error is reduced by factors up to 2.0 for rotations and 3.8 for translations, which is an enormous improvement.

Our work is valuable for stereovision. Extending it to the multi-view case is not trivial because of track consistency. First, removing one match does not necessarily remove the associated points from the track and leads to a substantially different bundle adjustment problem. Second, the location of points in a track would need to be optimized simultaneously in all associated images. We actually want *track selection* (or *reduction*) as well as *track refinement*. Besides, a good term to minimize to assess the benefit of match reduction is likely to be linked to the total reprojection error with respect to all 3D points *after* bundle adjustment. A study similar to that of Section 2 thus has to be carried out.

Still, a lower bound of the possible improvement can be obtained by applying match selection (MS) on each image pair in an SfM pipeline, before actual processing by the system. A preliminary experiment on Strecha et al.’s dataset using OpenMVG [21], a competitor to Bundler, shows improvements up to 15% on the average camera location error, in particular on scenes with wider viewpoint changes and less images (HerzJesu-P8 vs -P25, Castle-P19 vs -P30). Conversely, it may be the case that bundle adjustment is doing a good job at averaging on long tracks, compensating for the inaccuracy of point location. Track selection and track refinement are thus likely to be more profitable on difficult scenes.

Finally, most of our results are constructed on empirical studies. We however believe the “quality vs. quantity” issue deserves a better theoretical treatment, including a study of the influence of the configuration of points in images.

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References

1. Aanæs, H., Dahl, A.L., Pedersen, K.S.: Interesting interest points. *International Journal of Computer Vision* 97(1), 18–35 (2012)
2. Baatz, G., Köser, K., Chen, D., Grzeszczuk, R., Pollefeys, M.: Leveraging 3D city models for rotation invariant place-of-interest recognition. *IJCV* 96(3), 315–334 (2012), <http://dx.doi.org/10.1007/s11263-011-0458-7>
3. Baker, S., Scharstein, D., Lewis, J., Roth, S., Black, M.J., Szeliski, R.: A database and evaluation methodology for optical flow. *IJCV* 92(1), 1–31 (2011)
4. Brooks, M.J., Chojnacki, W., Gawley, D., Van Den Hengel, A.: What value covariance information in estimating vision parameters? In: *Proceedings of the 8th IEEE International Conference on Computer Vision (ICCV)*. vol. 1, pp. 302–308. IEEE (2001)
5. Cao, Y., McDonald, J.: Viewpoint invariant features from single images using 3D geometry. In: *WACV* (2009)
6. Choi, S., Kim, T., Yu, W.: Performance evaluation of RANSAC family. In: *BMVC*. pp. 1–12 (2009)
7. Chum, O., Matas, J.: Matching with PROSAC – progressive sample consensus. In: *CVPR* (2005)
8. Chum, O., Matas, J., Obdrzalek, S.: Enhancing RANSAC by generalized model optimization. In: *ACCV* (2004)
9. Fischler, M.A., Bolles, R.C.: Random sample consensus: a paradigm for model fitting with applications to image analysis and automated cartography. *CACM* 24(6) (1981)
10. Gruen, A.: Adaptive least squares correlation: a powerful image matching technique. *S. Afr. J. of Photogrammetry, Remote Sensing and Cartography* 14(3) (1985)
11. Hartley, R.I., Zisserman, A.: *Multiple View Geometry in Computer Vision*. Cambridge University Press (2004)
12. Kanazawa, Y., Kanatani, K.: Do we really have to consider covariance matrices for image features? In: *Proceedings of the 8th IEEE International Conference on Computer Vision (ICCV)*. vol. 2, pp. 301–306. IEEE (2001)
13. Köser, K., Koch, R.: Perspectively invariant normal features. In: *ICCV* (2007)
14. Köser, K., Koch, R.: Exploiting uncertainty propagation in gradient-based image registration. In: *BMVC* (2008)
15. Liu, Z., Marlet, R.: Virtual line descriptor and semi-local graph matching method for reliable feature correspondence. In: *BMVC* (2012)
16. Lowe, D.: Distinctive image features from scale-invariant keypoints. *IJCV* 60(2), 91–110 (2004)
17. Matas, J., Chum, O., Urban, M., Pajdla, T.: Robust wide-baseline stereo from maximally stable extremal regions. *Image and Vision Computing* 22(10), 761–767 (2004)
18. Mikolajczyk, K., Schmid, C.: Scale & affine invariant interest point detectors. *IJCV* 60(1), 63–86 (2004)
19. Moisan, L., Stival, B.: A probabilistic criterion to detect rigid point matches between two images and estimate the fundamental matrix. *IJCV* 57(3), 201–218 (2004)
20. Morel, J.M., Yu, G.: ASIFT: A new framework for fully affine invariant image comparison. *SIAM Journal on Imaging Sciences* 2(2), 438–469 (2009)
21. Moulon, P., Monasse, P., Marlet, R.: Adaptive Structure from Motion with a contrario model estimation. In: *ACCV* (2012)

22. Potůčková, M.: Image matching and its applications in photogrammetry. Ph.D. thesis, Aalborg Universitet (2004)
23. Strecha, C., von Hansen, W., Van Gool, L., Fua, P., Thoennessen, U.: On benchmarking camera calibration and multi-view stereo for high resolution imagery. In: CVPR (2008)
24. Sur, F., Noury, N., Berger, M.O.: Computing the uncertainty of the 8 point algorithm for fundamental matrix estimation. In: Proceedings of 19th British Machine Vision Conference (BMVC). pp. 96.1–96.10 (2008)
25. Tang, Z.: High precision in Camera calibration. Ph.D. thesis, ENS Cachan (2012)
26. Torr, P.H.S., Zisserman, A.: MLESAC: a new robust estimator with application to estimating image geometry. *CVIU* 78(1), 138–156 (2000)
27. Torr, P.H., Murray, D.W.: The development and comparison of robust methods for estimating the fundamental matrix. *IJCV* 24(3), 271–300 (1997)
28. Wu, C., Clipp, B., Li, X., Frahm, J.M., Pollefeys, M.: 3D model matching with viewpoint-invariant patches (VIP). In: CVPR (2008)
29. Zeisl, B., Georgel, P.F., Schweiger, F., Steinbach, E.G., Navab, N., Munich, G.: Estimation of location uncertainty for scale invariant features points. In: Proceedings of 20th British Machine Vision Conference (BMVC). pp. 1–12 (2009)